10/045,292

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FILE 'HOME' ENTERED AT 18:37:47 ON 26 DEC 2006

=> file reg

COST IN U.S. DOLLARS

SINCE FILE TOTAL ENTRY SESSION 0.21 0.21

FULL ESTIMATED COST

FILE 'REGISTRY' ENTERED AT 18:38:15 ON 26 DEC 2006
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STRUCTURE FILE UPDATES: 25 DEC 2006 HIGHEST RN 916309-42-7 DICTIONARY FILE UPDATES: 25 DEC 2006 HIGHEST RN 916309-42-7

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http://www.cas.org/ONLINE/UG/regprops.html

=> Uploading C:\Program Files\Stnexp\Queries\10045292c50.str

L1 STRUCTURE UPLOADED

=> d 11

L1 HAS NO ANSWERS

L1

claims 50251

Structure attributes must be viewed using STN Express query preparation.

=>

Uploading C:\Program Files\Stnexp\Queries\10045292c54.str

L2 STRUCTURE UPLOADED

=> d 12

L2 HAS NO ANSWERS

L2

Structure attributes must be viewed using STN Express query preparation.

=> Uploading C:\Program Files\Stnexp\Queries\10045292c55.str

L3 STRUCTURE UPLOADED

L3 HAS NO ANSWERS
L3 STR

Structure attributes must be viewed using STN Express query preparation.

=>
Uploading C:\Program Files\Stnexp\Queries\10045292c56.str

L4 STRUCTURE UPLOADED

=> d 14 L4 HAS NO ANSWERS L4 STR

Cl. Ul & Sle

Structure attributes must be viewed using STN Express query preparation.

Uploading C:\Program Files\Stnexp\Queries\10045292c57.str

L5 STRUCTURE UPLOADED

STR

=> d 15

L5 HAS NO ANSWERS

L5

Structure attributes must be viewed using STN Express query preparation.

=> s 11

SAMPLE SEARCH INITIATED 18:39:57 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 2885 TO ITERATE

69.3% PROCESSED 2000 ITERATIONS INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

24 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
BATCH **COMPLETE**

PROJECTED ITERATIONS: 54479 TO 60921 PROJECTED ANSWERS: 339 TO 1045

L6 24 SEA SSS SAM L1 .

=> s 12

SAMPLE SEARCH INITIATED 18:40:00 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 959 TO ITERATE

100.0% PROCESSED 959 ITERATIONS 1 ANSWERS SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED ITERATIONS: 17323 TO 21037

PROJECTED ANSWERS: 1 TO 80

L7 1 SEA SSS SAM L2

=> s 13

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SAMPLE SCREEN SEARCH COMPLETED - 34 TO ITERATE

100.0% PROCESSED 34 ITERATIONS 2 ANSWERS SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**

BATCH **COMPLETE**

PROJECTED ITERATIONS: 331 TO 1029

PROJECTED ANSWERS: 2 TO 124

L8 2 SEA SSS SAM L3

=> s 14

SAMPLE SEARCH INITIATED 18:40:05 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED - 46 TO ITERATE

100.0% PROCESSED SEARCH TIME: 00.00.01

46 ITERATIONS

O ANSWERS

FULL FILE PROJECTIONS: ONLINE **COMPLETE** BATCH **COMPLETE**

PROJECTED ITERATIONS:

PROJECTED ANSWERS:

514 TO 1326 0 TO

O SEA SSS SAM L4

=> s 15

L9

SAMPLE SEARCH INITIATED 18:40:08 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED -40 TO ITERATE

100.0% PROCESSED

40 ITERATIONS

1 ANSWERS

SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE** BATCH **COMPLETE**

PROJECTED ITERATIONS:

421 TO 1179

PROJECTED ANSWERS:

1 TO 80

L10

1 SEA SSS SAM L5

=> file caplus

COST IN U.S. DOLLARS

FULL ESTIMATED COST

SINCE FILE TOTAL

ENTRY

1.76

SESSION 1.97

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FILE COVERS 1907 - 26 Dec 2006 VOL 146 ISS 1 FILE LAST UPDATED: 25 Dec 2006 (20061225/ED)

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=> s 16

L111835 L6

=> s 17

L12 2 L7

=> s 18

L13 60 L8

=> s 19L14

0 L9

=> s 110

L15 1 L10

=> d bib abs hitstr 1-2 112

L12 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2006 ACS on STN

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10/908,624
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AN 1968:3159 CAPLUS DN 68:3159 ΤI Pyrimidine nucleosides Merck and Co., Inc. PA Neth. Appl., 21 pp. CODEN: NAXXAN DT Patent LA Dutch FAN.CNT 1 PATENT NO. KIND DATE DATE APPLICATION NO. PΙ NL 6614805 19670425 NL 1966-14805 19661020 DE 1620047 DE FR 1502956 FR GB 1117039 GB US 3346561 19651024 19671010 US 1965-505029 PRAI US 19651024 AB The preparation of the α - and β -anomers of 1-(3-deoxy-Dribofuranosyl)pyrimidines by treating a 3-deoxy-D-ribofuranosyl halide with a 2,4-dialkoxypyrimidine and subsequent solvolysis of the product obtained is described. Thus, a solution of 2,5-di-0-benzoyl-3-deoxy- β -Dribofuranosyl bromide (prepared from 2 g. methyl 2,5/di-O-benzoyl-3-deoxy- β -D-ribofuranoside) in 20 ml. anhydrous CH2Cl2 was freated with stirring with a solution of 1.9 g. 2,4-dimethoxy-5-fluoropyrimidine in 80 ml. anhydrous CH2Cl2 for 80 hrs. at 25° to give 39% 1-(2,5-di-0,benzoyl-3-deoxy- β -D-ribofuranosyl)-5-fluoro-4-methoxy-2(1H)-pyrimidinone (I), m. 148-50°, [α] 578 27°, [α] D 24° (c 0.64, CHCl3). A solution of 234 mg. I in 5 ml. MeOH and 0.6 ml. 2.5N NaOH was heated 1.5 hrs. at 60° and evaporated to dryness, the residue (428 mg.) dissolved in 10 ml. H2O, and the solution treated with 1 g. Dowex 50 (H+) resin and stirred 10 min. to yield 65% 5-flugro-3'-deoxyuridine, m. $166.5-7.5^{\circ}$, [\alpha]D 30\circ, [\alpha]578 33\circ (c 1.1, H2O). A mixture of 94 mg. I and 1.4 ml. Et/OH saturated at 0° with NH3 was heated in a closed tube 12 hrs. at 100% to yield 68% 3'-deoxy-5-fluorocytidine hydrogen sulfate, m. 175-6°, $[\alpha]D$ 41°, $[\alpha]$ 578 45° (c 0.51, H2O). Similar reaction of 4.47 g. 2,4-dimethoxy-5-methylpyrimidine/and 6.67 g. 2,5-di-O-pnitrobenzoyl-3-deoxy- β -D-ribofuranosyl β romide gave 420 mg. pure 1-(2,5-di-0-p-nitrobenzoyl-3-deoxy- β -D-kibofuranosyl)-4-methoxy-5methyl-2(1-H)-pyrimidinone (II), m. 163-7° (EtOAc, petr. ether). Further elution of the column with CHC/13 gave 500 mg. II, m. 217-18° which when recrystd. from a mixture of CHCl3 and MeOH gave 440 mg. II, m. 218-19°. Repeated crystallization of the α -rich fractions gave 710 mg. α -D-anomer, m/ 218-20°. A total of 18% α - and 28% β -II was obtained. A mixture of 400 mg. II and 5 ml. MeOH, saturated with NH3 at 0° was heated 16 hrs. at 100° and worked up to yield 52% 1-(3-de ϕ xy- β -D-ribofuranosyl)-5methylcytosine, m. 223-6°, $[\alpha]D$ 30 $[\alpha]$ 578 32° (c 0.59, H2O). A suspension df 960 mg. II in 21 ml. anhydrous MeOH was treated with 60 mg. Na in 3 ml. MeOH, the solution refluxed 1.25 hrs., evaporated to dryness, 30 ml. H2O added and further worked up with Et2O to yield 66% 1-(3-deoxy-β-D-ribofuranosyl)-4-methoxy-5-methyl-2(1H)pyrimidinone (III), m. $185-7^{\circ}$ [α] D -157° , [α] 578 -166° (c 0.99, H2O). A solution of 1.54 g. II in 34 ml. anhydrous MeOH was treated with 100 mg. Na in 3 ml. anhydrous MeOH, the mixture refluxed 1 hr., solvent removed, the residue treated with H2O, filtered, washed with H2O, and the filtrate treated with 15 g. Dowex 50W (H+) resin and worked up to yield 258 mg. III, m. $196-8^{\circ}$ (MeOH), $[\alpha]$ 278 27° (c 0.765, H2O). A solution of 279 mg. III in 10 ml. anhydrous MeOH was treated with 1 ml. 30% HCl in MeOH and the solution kept 6 days at 25° and evaporated to dryness to yield 76% 1-(3-deoxy- α -D-ribofuranosyl)-5methyluracil (IV), m. $188-91^{\circ}$ (MeOH), $[\alpha]D-112^{\circ}$, [α] $5\overline{7}8$ -118° (c 0.17, H2O). A similar treatment of III, m. 196-8°, gave 81% IV m. 96-100°, solidified and m. 155-7°, $[\alpha]D$ 1.4°, $[\alpha]$ 578 2.3 (c 0.44, H2O). Also prepared was 11% 1-(2,5-di-O-p-nitrobenzoyl-3-deoxy- β -Dribofuranosyl)-4-methoxy-2(1H)-pyrimidinone (V), m. 193-4° (EtOAc-petroleum ether), $[\alpha]D - 9.2^{\circ}$ (c 1.09, CHCl3). Treatment of 760 mg. V in 17 ml. anhydrous MeOH with 60 mg. Na in 3 ml. anhydrous MeOH as described above gave 46% 1-(3-deoxy- α -Dribofuranosyl)-4-methoxy 2(1H)-pyrimidinone (VI), m. 209-11°, $[\alpha]D -182^{\circ}$, $[\alpha]578 -194^{\circ}$ (c 0 263, H2O). Also prepared was 75% 1-(3-deoxy-β-D-ribofuranosyl)-4-methoxy-2(1H)pyrimidinone, m. 187-91°. Reaction of 130 mg. VI in 5 ml. MeOH with 0.5 ml. 31% HCl in MeOH gave 27% 1-(3-deoxy- α -D-

Ch. zarsy backrimi Mear backrims

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McIntosh

ribofuranosyl)uracil m. 125.5-6.5° (MeOH-Et2O), $[\alpha]D$ -134° , [α] 578 -141° (c 0.134, H2O). A solution of 300 mg. V in 4 ml. MeOH saturated with NH3 at 0° was heated overnight at 100°, evaporated to dryness, 20 ml./H2O added, the mixture filtered and worked up to yield 82% 1-(3-deoxy- α -D-ribofuranosyl)cytosine, m. 225-9°, $[\alpha]D = 130°$, $[\alpha]578 = 141°/(c 0.73)$ H2O). Also prepared was 80% $1-\sqrt{3}$ -deoxy- β -D-ribofuranosyl)cysotine, m. 224-30°, [α]D 54°, [α]578 58° $\langle c$ 0.71, H2O). 7139-62-0P ITRL: SPN (Synthetic preparation); PREP (Preparation) (preparation of) RN 7139-62-0 CAPLUS Cytosine, 1-(3-deoxy- α /D-erythro-pentofuranosyl)- (7CI, 8CI) CN INDEX NAME) Absolute stereochemistry HO L12 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2006 ACS on STN 1966:93783 CAPLUS AN 64:93783 DN OREF 64:17699c-d 3'-Deoxynucleosides. IV. Pyrimidine 3'-deoxynucleosides Walton, Edward; Holly, Frederick W.; Boxer, George E.; Nutt, Ruth F. AU Merck Sharp & Dohme Res. Lab., Rahway, NJ CS Journal of Organic Chemi/stry (1966), 31(4), 1163-9 CODEN: JOCEAH; ISSN: 00/22-3263 Journal DT English LAcf. CA 63, 18243a. The 1-(3-deoxy-D-erythro-pentofuranosyl) derivs. of uracil, cytosine, thymine, and 5-methylcytosine were synthesized via Hilbert-Johnson reactions of 2,5-di-O-acyl-3-deoxy-D-erythropentofuranosyl bromide with the appropriate 2,4-dialkoxypyrimidine followed by methanolysis or ammonolysis. Both anomers were produced although the traps rule predicts that only trans (β) coupling products would be formed. The optical rotations of the anomeric pairs were found to be the reverse of those predicted by Hudson's rules of isorotation. Assignments of the anomeric configuration of the products and intermediates were made, in part, on the basis of N.M.R. as well as optical rotatory dispersion. Some properties characteristic of the anomers are tabulated. 7139-62-0P,/Cytosine, 1-(3-deoxy- α -D-erythro-pentofuranosyl)-ITRL: PREP (Preparation) (preparation of) 7139-62-0/ CAPLUS RNCytosine, $\int 1-(3-\text{deoxy}-\alpha-D-\text{erythro-pentofuranosyl}) - (7CI, 8CI)$ (CA CN INDEX NAME) Jedy Winger La Absolute stereochemistry. H₂N

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10/908,624
=> d bib abs hitstr 1 115
L15 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2006 ACS on STN
   1995:938831 CAPLUS
AN
   124:117872
DN
    Syntheses of [6,7-15N]-Adenosine, [6,7-15N]-2'-Deoxyadenosine, and
ΤI
    [7-15N]-Hypoxanthine
    Pagano, Alex R.; Lajewski, Wayne M.; Jones, Roger A.
AU
    Department of Chemistry, Rutgers, State University of New Jersey,
    Piscataway, NJ, 08855, USA
    Journal of the American Chemical Society (1995), 117(47), 11669-72
SO
    CODEN: JACSAT; ISSN: 0002-7863
PB
    American Chemical Society
DT
    Journal
LA
    English
    We have developed a high-yield foute for the synthesis of
     [7-15N]-hypoxanthine in four steps in an overall yield of 81%. This
    procedure uses [15N]-sodium nitrite as the 15N source and an inexpensive
    pyrimidine to provide an económical route to this useful 15N-labeled
    intermediate. Conversion to (7-15N)-6-chloropurine followed by enzymic
    trans-glycosidation gives the corresponding ribo- and 2'-
    deoxyribonucleosides. Ammonolysis of the 6-chloro moiety to give the
    [6,7-15N]-labeled nucleosides is effected simply and in high yield using 2
    equiv of [15N]-ammonium chloride and 3 equiv of potassium bicarbonate.
   173170-82-6P
    RL: BPN (Biosynthetic preparation); RCT (Reactant); BIOL (Biological
    study); PREP (Preparation); RACT (Reactant or reagent)
        (syntheses of N-14 Vabeled nucleosides with labeled sodium nitrite)
RN
    173170-82-6 CAPLUS
    9H-Purine-7-15N, 6-ch/oro-9-β-D-ribofuranosyl- (9CI) (CA INDEX NAME)
CN
Absolute stereochemistry.
  Cl
       15N
                           OH
                R
                 R S
             HO
                      OH
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     (FILE 'HOME' ENTERED AT 18:37:47 ON 26 DEC 2006)
     FILE 'REGISTRY' ENTERED AT 18:38:15 ON 26 DEC 2006
L1
                STRUCTURE UPLOADED
L2
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L5
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L11
           1835 S L6
L12
              2 S L7
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11641 HCV

=> s 116 and hcv

=> s 111 or 113

60 S L8

0 S L9

1 S L10

1891 L11 OR L13

L13

L14

L15

L16

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22 HCVS
         11645 HCV
                 (HCV OR HCVS)
L17
            24 L16 AND HCV
=> d bib abs hitstr 1-24 117
    ANSWER 1 OF 24 CAPLUS
                             COPYRIGHT 2006 ACS on STN
AN
     2005:1261770 CAPLUS
    144:7097
DN
TI
     Preparation of macrocyclic carboxylic acid derivatives as inhibitors of
    HCV replication
    Blatt, Lawrence M.; Andrews, Steven W.; Condroski, Kevin R.; Doherty,
IN
    George A.; Jiang, Yutong; Josey, John A.; Kennedy, April L.; Madduru,
    Machender R.; Stengel, Peter J.; Wenglowsky, Steven M.; Woodard, Benjamin
    T.; Woodard, Laura
    USA
PA
SO
    U.S. Pat. Appl. Publ., 228 pp., Cont.-in-part of U.S. Ser. No. 64,445.
    CODEN: USXXCO
DT
     Patent
     English
LA
FAN.CNT 3
     PATENT NO.
                         KIND
                                 DATE
                                             APPLICATION NO.
                                                                    DATE
     US 2005267018
PΙ
                          A1
                                 20051/201
                                             US 2005-93884
                                                                    20050329
                                 20050428
    WO 2005037214
                          A2
                                             WO 2004-US33970
                                                                    20041013
    WO 2005037214
                          A3
                                 20051103
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             CN, CO, CR, CU, CZ, DE,
                                     DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD,
             GE, GH, GM, HR, HU, ID/
                                     IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC,
             LK, LR, LS, LT, LU, LV,
                                     MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI,
             NO, NZ, OM, PG, PH, PL,
                                     PT, RO, RU, SC, SD, SE, SG, SK, SL, SY,
                                 TZ, UA, UG, UZ, VC, VN, YU, ZA, ZM, ZW
             TJ, TM, TN, TR, TT,
         RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM,
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* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

20031014

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20040414

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20040922

20041013

20050223

SN, TD, TG

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A1

PRAI US 2003-511541P

US 2004-558161P

US 2004-562418P

US 2004-612381P

US 2004-612460P

WO 2004-US33970

MARPAT 144:7097

US 2005-64445

OS GI AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE,

AB The invention relates to macrocyclic compds., e.g., I [Q is (un) substituted 2-isoindolinyl, 2-isoquinolinyl, 1-benzoazetidinyl, 1-indolinyl, (3,4-dehydro)pyrrolidino, (3,4-dehydro)piperidino or Q is R3-R4, where R3 is alkyl, cycloalkyl, alkylcycloalkyl, Ph, pyridyl and other heterocyclic groups and R4 is H, Ph, pyridyl and other heterocyclic groups; V is O, S, NH; W is O, NR5 or CR5, where R5 is H, alkyl, fluoroalkyl, cycloalkyl, alkylcycloalkyl; Y is a sulfonimide CONHSO2R6, where R6 is (un) substituted alkyl, fluoroalkyl, cycloalkyl, alkylcycloalkyl, aryl, heteroaryl or (un) substituted phenyl; or Y is carboxy or a pharmaceutically-acceptable salt or prodrug; R1 is H, (un) substituted alkyl, cycloalkyl, alkylcycloalkyl, Ph or benzyl; R2 is H, alkyl, (thio)carbamoyl, acyl, or sulfonyl group; the dashed line represents an optional double bond], for use in pharmaceutical compns. for the treatment of hepatitis C virus (HCV) infection and liver fibrosis. Thus, compound II, prepared by reaction of the macrocyclic prolinol derivative with CDI and 4-fluoro-2,3-dihydro-1H-isoindole, showed IC50 and EC50 $< 0.1 \mu M$ in the NS3-NS4A protease inhibition assay and did not display toxicity in Rattus sp. when dosed for seven days at 30 mg/kg BID, providing at least a 10-fold safety margin above the presumptive efficacious dose (3 mg/kg) that yields liver concns. 100-fold in excess of the replicon EC50 value of the compound 7481-89-2 IT

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (preparation of macrocyclic carboxylic acid derivs. as inhibitors of HCV replication)

RN 7481-89-2 CAPLUS

Cytidine, 2',3'-dideoxy- (8CI, 9CI) (CA INDEX NAME) CN

Absolute stereochemistry. Rotation (+).

L17 ANSWER 2 OF 24 CAPLUS COPYRIGHT 2006 ACS on STN

2005:1151410 CAPLUS AN

145:336253 DN

Synthesis and in vitro anti-HCV activity of β -D- and L-2'-deoxy-2'-fluororibonucleosides

Shi, Junxing; Du, Jinfa; Ma, Tianwei; Pankiewicz, Krzysztof W.; Patterson, Steven E.; Hassan, Abdalla E. A.; Tharnish, Phillip M.; McBrayer, Tamara R.; Lostia, Stefania; Stuyver, Lieven J.; Watanabe, Kyoichi A.; Chu, Chung K.; Schinazi, Raymond F.

CS Pharmasset, Inc., Tucker, GA, USA

Nucleosides, Nucleotides & Nucleic Acids (2005), 24(5-7), 875-879 CODEN: NNNAFY; ISSN: 1525-7770

PBTaylor & Francis, Inc.

DTJournal

LA English

OS

CASREACT 145:336253 Based on the discovery of $\beta\text{-D-2'-deoxy-2'-fluorocytidine}$ as a potent anti-hepatitis C virus (HCV) agent, a series of $\beta\text{-}D\text{-}$ and L-2'-deoxy-2'-fluororibonucleosides with modifications at 5 and/or 4 positions were synthesized and evaluated for their in vitro activity against HCV and bovine viral diarrhea virus (BVDV). The introduction of the 2'-fluoro group was achieved by either fluorination of 2,2'-anhydronucleosides with hydrogen fluoride-pyridine or potassium fluoride, or a fluorination of arabinonucleosides with DAST. Among the analogs synthesized, only the 5-fluoro compds., namely β -D-2'-deoxy-2',5-difluorocytidine, had anti- HCV activity in the subgenomic HCV replicon cell line, and inhibitory activity against rRNA. As β -D-N4-hydroxycytidine (NHC) had previously shown potent anti-HCV activity, the two functionalities of the N4-hydroxyl and the 2'-fluoro were combined into one mol., yielding β -D-2'-deoxy-2'-fluoro-N4-hydroxycytidine. However, this nucleoside showed neither anti-HCV activity nor toxicity. All the L-forms of the analogs were devoid of anti-HCV activity. None of the compds. showed anti-BVDV activity, suggesting that the BVDV system cannot reliably predict anti-HCV activity in vitro.

3258-02-4 IT

> RL: PAC (Pharmacological activity); BIOL (Biological study) (preparation and anti-HCV, anti-BVDV, rRNA inhibition activity of β-D- and L-2'-deoxy-2'-fluororibonucleosides via fluorination of anhydronucleosides and arabinonucleosides)

3258-02-4 CAPLUS RN

Uridine, 4-oxime (9CI) (CA INDEX NAME) CN

Absolute stereochemistry.

RE.CNT 14 THERE ARE 14 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L17 ANSWER 3 OF 24 CAPLUS COPYRIGHT 2006 ACS on STN
AN 2005:1106860 CAPLUS
DN 143:367596

TI Preparation of macrocyclic carboxylic acids or sulfonamides as inhibitors of HCV replication

IN Blatt, Lawrence M.; Wenglowsky, Steven M.; Andrews, Steven W.; Condroski, Kevin R.; Jiang, Yutong; Kennedy, April L.; Doherty, George A.; Josey, John A.; Stengel, Peter J.; Woodard, Benjamin T.; Madduru, Machender R.

PA Intermune, Inc., USA SO PCT Int. Appl., 444 pp. CODEN: PIXXD2

DT Patent

FAN.CNT 3																		
LAN.	PATENT NO.				KIND		DATE /		•	APPLICATION NO.				DATE				
PI	WO 2005095403 WO 2005095403			A2 A3			1013 1201	1	WO 2005-US10494					20050329				
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	AU 2005	-	-		A1		2005	1013		AU 2	005-	2288	94		2	0050	329	
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	US 2004	-5624	118P		P		2004	0414										
	US 2004	-6123	381P		P		200h	0922										
	US 2004				P		200A	0922										
	WO 2005				W		2005	0329										
os	MARPAT :	143:3	36759	96														
GI							l	/										

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

The invention relates to macrocyclic compds., e.g., I [Q is AB (un) substituted 2-isoindolinyl, 2-isoquinolinyl, 1-benzoazetidinyl, 1-indolinyl, (3,4-dehydro)pyrrolidino, (3,4-dehydro)piperidino or Q is R3-R4, where R3 is alkyl, cycloalkyl, alkylcycloalkyl, Ph, pyridyl and other heterocyclic groups and R4 is H, Ph, pyridyl and other heterocyclic groups; V is O, S, NH; W is O, NR5 or CR5, where R5 is H, alkyl, fluoroalkyl, cycloalkyl, alkylcycloalkyl; Y is a sulfonimide CONHSO2R6, where R6 is (un) substituted alkyl, fluoroalkyl, cycloalkyl, alkylcycloalkyl, aryl, heteroaryl or (un)substituted phenyl; or Y is carboxy or a pharmaceutically-acceptable salt or prodrug; R1 is H, (un) substituted alkyl, cycloalkyl, alkylcycloalkyl, Ph or benzyl; R2 is H, alkyl, (thio)carbamoyl, acyl, or sulfonyl group; the dashed line represents an optional double bond], for use in pharmaceutical compns. for the treatment of flaviviral or hepatitis C virus (HCV) infection and liver fibrosis. Thus, compound II, prepared by reaction of the macrocyclic prolinol derivative with CDI and 4-fluoro-2,3-dihydro-1Hisoindole, showed IC50 and EC50 < 0.1 μM in the NS3-NS4A protease inhibition assay and did not display toxicity in Rattus sp. when dosed for seven days at 30 mg/kg BID, providing at least a 10-fold safety margin above the presumptive efficacious dose (3 mg/kg) that yields liver concns. 100-fold in excess of the replicon EC50 value of the compound IT7481-89-2

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (preparation of macrocyclic carboxylic acids or sulfonamides as inhibitors of HCV replication)

RN 7481-89-2 CAPLUS

CN Cytidine, 2',3'-dideoxy- (8CI, 9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

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ANSWER 4 OF 24 CAPLUS
                             COPYRIGHT 2006 ACS on STN
     2005:371064 CAPLUS
AN
     142:430532
DN
     Preparation of macrocyclic carboxylic acids and acylsulfonamides as
ΤI
     inhibitors of HCV replication
     Blatt, Lawrence M.; Wenglowsky, Steven Mark; Andrews, Steven Wade; Jiang,
ΙN
     Yutong; Kennedy, April Layne; Condroski, Kevin Ronald; Josey, John
     Anthony; Stengel, Peter John; Madduru, Machender R.; Doherty, George
     Andrew; Woodard, Benjamin T.
PA
     Intermune, Inc., USA; Array Biopharma Inc.
SO
     PCT Int. Appl., 244 pp.
     CODEN: PIXXD2
     Patent
\mathsf{DT}
LA
     English
FAN.CNT 3
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                                             APPLICATION NO.
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PRAI US 2003-511541P
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     US 2005-64445
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OS
    MARPAT 142:430532
GI
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- * STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY AVAILABLE VIA OFFLINE PRINT *
- The invention relates to macrocyclic compds., e.g., tetrahydroisoquinolinecarboxylic acid derivs. I [R1, R2 are independently H, halo, cyano, hydroxy, alkyl, alkoxy; R5 is a carbamoyl, acyl or carboxy ester; Y is a sulfonimide CONHSO2R9, where R9 is alkyl, cycloalkyl or (un)substituted phenyl; or Y is carboxylic acid or pharmaceutically-acceptable salt or prodrug; R10, R11 are independently H or alkyl or CR10R11 is cyclopropyl, cyclobutyl, cyclopentyl or cyclohexyl; W is O or NH; the dashed line represents an optional double bond], for use in pharmaceutical compns. for the treatment of hepatitis C virus (HCV) infection and liver fibrosis. Thus, compound II, prepared by reaction of the macrocyclic prolinol derivative with CDI and 4-fluoro-2,3-dihydro-1H-

isoindole, showed IC50 and EC50 < 0.1 μM in the NS3-NS4A protease inhibition assay and did not display toxicity in Rattus sp. when dosed for seven days at 30 mg/kg BID, providing at least a 10-fold safety margin above the presumptive efficacious dose (3 mg/kg) that yields liver concns. 100-fold in excess of the replicon EC50 value of the compound

IT 7481-89-2, 2' 3' Dideoxycytidine

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (preparation of macrocyclic carboxylic acids and acylsulfonamides as inhibitors of HCV replication)

RN 7481-89-2 CAPLUS

CN Cytidine, 2',3'-dideoxy- (8CI, 9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

L17 ANSWER 5 OF 24 CAPLUS COPYRIGHT 2006 ACS on STN

KIND

AN 2005:185375 CAPLUS

DN 142:254563

TI Antimetabolite antiviral dosing regimen for hepatitis C virus or flaviviridae therapy

DATE

IN Stuyver, Lieven J.

PA Belg.

SO U.S. Pat. Appl. Publ., 23 pp.

CODEN: USXXCO

PATENT NO.

DT Patent

LA English

FAN.CNT 2

	~				
ΡI	US 2005049220	A1	20050 3 03	US 2004-921052	20040818
PRAI	US 2003-496202P	P	20039⁄818		
AB	_	-		antimetabolite to the	
	cannot be administed	red on	a daily or	chronic basis as is us	ual in
	antiviral therapy (referre	d to below	as an "anti-HCV	
				using a traditional a	
	dosing regimen (for	exampl	e via i.v.	or parenteral injection	n), over a
				ion of therapy until re	
	viral load is noted	. This	dosing reg	imen runs counter to co	onventional
	antiviral experience	e, wher	ein effecti	ve agents are usually a	administered

APPLICATION NO.

DATE

indefinite daily basis. IT 7481-89-2, Zalcitabine

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(antimetabolite antiviral dosing regimen for hepatitis C virus or flaviviridae therapy)

over at least fourteen days of sustained therapy, and typically on an

RN 7481-89-2 CAPLUS

CN Cytidine, 2',3'-dideoxy- (8CI, 9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

L17 ANSWER 6 OF 24 CAPLUS COPYRIGHT 2006 ACS on STN

AN 2005:177803 CAPLUS

DN 142:254560

TI Antimetabolite antiviral dosing regimen for hepatitis C virus or flaviviridae therapy

```
IN
     Stuyver, Lieven J.
     Pharmasset, Inc., USA
PA
     PCT Int. Appl., 61 pp.
     CODEN: PIXXD2
DT
     Patent
     English
LA
FAN.CNT 2
     PATENT NO.
                         KIND
                                             APPLICATION NO.
                                DATE
                                                                    DATE
PI
                                20050/303
     WO 2005018330
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                                            WO 2004-US26686
                                                                    20040817
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             LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI,
             NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY,
             TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
         RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM,
            AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK,
             EE, ES, FI, FR, GB, $R, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE,
             SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE,
             SN, TD, TG
PRAI US 2003-496202P
                          Р
                                20030818
    An anti-hepatitis C agent which is an anti-metabolite to the host and
    cannot be administered on a daily or chronic basis as is usual in
    anti-viral therapy (referred to below as an "anti-HCV
    anti-metabolite"), can be administered using a traditional anti-cancer
     dosing regimen (for example via i.v. or parenteral injection), over a
    period of 1-7 days followed by cessation of therapy until rebound of the
    viral load is noted. This dosing regimen runs counter to conventional
    antiviral experience, wherein effective agents are usually administered
    over at least fourteen days of sustained therapy, and typically on an
     indefinite daily basis.
IT
    7481-89-2, Zalcitabine
    RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL
     (Biological study); USES (Uses)
        (antimetabolite antiviral dosing regimen for hepatitis C virus or
        flaviviridae therapy)
     7481-89-2 CAPLUS
RN
    Cytidine, 2',3'-dideoxy- (8CI, 9CI) (CA INDEX NAME)
CN
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Absolute stereochemistry. Rotation (+).

RE.CNT 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

```
L17 ANSWER 7 OF 24 CAPLUS COPYRIGHT 2006 ACS on STN
    2004:1065490 CAPLUS
AN
DN
    142:147801
    Metabolism of the anti-hepatitis C virus nucleoside \beta-D-N4-
    hydroxycytidine in different liver cells
    Hernandez-Santiago, Brenda I.; Beltran, Thierry; Stuyver, Lieven; Chu,
    Chung K.; Schinazi, Raymond F.
     Department of Pediatrics, Emory School of Medicine, Decatur, USA
    Antimicrobial Agents and Chemotherapy (2004), 48(12), 4636-4642
     CODEN: AMACCQ; ISSN: 0066-4804
PB
    American Society for Microbiology
DT
    Journal
LA
     English
     \beta-D-N4-Hydroxycytidine (NHC) was found to have selective
     anti-hepatitis C virus (HCV) activity in the HCV
     replicon system (clone A). The intracellular metabolism of tritiated NHC was
     investigated in the HCV replicon system, Huh-7 cells, HepG2
     cells, and primary human hepatocytes. Incubation of cells with 10 µM
     radiolabeled NHC demonstrated extensive and rapid phosphorylation in all
     liver cells. Besides the 5'-mono, -di-, and -triphosphate metabolites of
     NHC, other metabolites were characterized. These included cytidine and
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uridine mono-, di-, and triphosphates. UTP was the predominant early metabolite in Huh-7 cells and primary human hepatocytes, suggesting deamination of NHC as the primary catabolic pathway. The intracellular half-lives of radiolabeled NHC-triphosphate and of CTP and UTP derived from NHC incubation in Huh-7 cells were calculated to be 3.0 ± 1.3 , 10.4±3.3, and 13.2±3.5 h, resp. Studies using monkey and human whole blood demonstrated more-rapid deamination and oxidation in monkey cells than in human cells, suggesting that NHC may not persist long enough in plasma to be delivered to liver cells.

3258-02-4

RL: PKT (Pharmacokinetics); BIOL (Biological study) (metabolism of the anti-hepatitis C virus nucleoside $\beta\text{-}D\text{-}N4\text{-}$ hydroxycytidine in different liver cells)

3258-02-4 CAPLUS RN

CN Uridine, 4-oxime (9CI) (CA INDEX NAME)

Absolute stereochemistry.

THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 20 ALL CITATIONS AVAILABLE IN THE RE FORMAT

L17 ANSWER 8 OF 24 CAPLUS COPYRIGHT 2006 ACS on STN

2004:703121 CAPLUS AN

141:207236 DN

Preparation of 1,1-dioxido-4H-1,2,4-benzothiadiazines as hepatitis C polymerase inhibitors and anti-infective agents

Pratt, John K.; Betebenner, David A.; Donner, Pamela L.; Green, Brian E.; Kempf, Dale J.; McDaniel, Keith F.; Maring, Clarence J.; Stoll, Vincent S.; Zhang, Rong

PA USA

SO U.S. Pat. Appl. Publ., 278 pp.

CODEN: USXXCO

DT Patent

FAN CNT 1

twn.	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
	US 2004167123 US 2002-423209P US 2003-461784P US 2003-489448P US 2003-509107P	A1 P P P	20040826 20021101 20030410 20030723 20031006	US 2003-699513	20031031
OS GI	MARPAT 141:207236				

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Title compds. I [wherein A = monocyclic or bicyclic ring selected from hetero/aryl, cycloalkyl, cycloalkenyl, heterocyclyl; R1 = H, (un) substituted cycloalkyl/cyclo/alkenyl, alkoxycarbonyl/alkoxy/aryl/aryls ulfonyl/arylsulfanyl/carboxy/cyano/heteroaryl/alkyl, heterocyclyl, etc.; R2, R3 = independently H, cyano, halo, (un) substituted alkenyl, alkoxycarbonyl, alkyl, heteroaryl, etc.; CR2R3C = 5- or 6-membered ring selected from Ph, pyridinyl, pyrimidinyl, pyridazinyl, thienyl, furanyl, pyrazolyl, oxazolyl, thiazolyl, imidazolyl, isoxazolyl, isothiazolyl, triazolyl, thiadiazolyl, tetrazolyl, cyclopentyl, and cyclohexyl; R4 = OH and derivs., halo, NH2 and derivs., etc.; R5 = independently CN, NO2, (un) substituted alk(en/yn)yl, hetero/aryl, arylsulfonyl, heterocyclyl etc.; n = 0-4; their pharmaceutically acceptable salts, stereoisomers, or tautomers] were prepared as hepatitis C (HCV) polymerase inhibitors for treating related infections. Thus II was prepared by

RN

CN

alkylation of III (preparation given) with tris(methylthio)methyl Me sulfate in AcOH, cyclization with 2-amino-4[(4-methoxymethoxy)methyl]thiophene-3-sulfonamide, deprotection, condensation with cyclopropanecarboxaldehyde, reduction with LiBH4. I inhibited HCV polymerase with IC50's in the range of 0.002 μM to 500 μM. I inhibited RNA replication with EC50 in the range of 0.002 μM to > 100 μM. I exhibited a cytopathic effect reduction with TC50's in the range of 6.6 μM to > 100 μM. 7481-89-2, Zalcitabine RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (combination therapy; preparation of 1,1-dioxidobenzothiadiazines as hepatitis C polymerase inhibitors and anti-infective agents) 7481-89-2 CAPLUS Cytidine, 2',3'-dideoxy- (8CI, 9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

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L17 ANSWER 9 OF 24 CAPLUS COPYRIGHT 2006 ACS on STN
     2004:453332 CAPLUS
AN
    141:17577
DN
     Concurrent inhibiting viral replication and treating cancer by pegylated
     arginine deiminase, and methods for determining the sensitivity to
     arginine deprivation therapy
IN
    Clark, Mike A.
PA
     Phoenix Pharmacologics, Inc., USA
SO
     PCT Int. Appl., 89 pp.
     CODEN: PIXXD2
DT
     Patent
LA
     English
FAN.CNT 1
     PATENT NO.
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                                DATE
                                             APPLICATION NO.
                                                                    DATE
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             GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK,
             LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ,
             OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM,
             TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
         RW: GH, GM, KE, LS, MW, MZ;
                                     SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,
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             FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR,
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PRAI US 2002-427497P
                                 20021118
                          Р
     WO 2003-US30770
                          W
                                 20030929
     The present invention is directed to methods of modulating viral
AB
     replication comprising administering to a patient arginine deiminase (ADI)
     bonded to polyethylene glycol (PEG). The present invention is also
     directed to methods of concurrently modulating viral replication and
     treating cancer, including, for example, sarcomas, hepatomas and
     melanomas. The present invention is also directed to methods of determining the
     susceptibility of an individual to arginine deprivation therapy for a
     viral infection, methods for improving liver function, and the like.
IT
     7481-89-2, Zalcitabine
     RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (dideoxycytosine, co-treatment with; concurrent inhibiting viral
        replication and treating cancer by pegylated arginine deiminase, and
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methods for determining sensitivity to arginine deprivation therapy) RN 7481-89-2 CAPLUS CN Cytidine, 2',3'-dideoxy- (8CI, 9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

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L17
    ANSWER 10 OF 24 CAPLUS COPYRIGHT 2006 ACS on STN
AN
     2004:412943 CAPLUS
     140:423711
DN
     Preparation of 1,1-dioxido-4H-1,2,4-benzothiadiazines as hepatitis C
     polymerase inhibitors and anti-infective agents
    Pratt, John K.; Betebenner, David A.; Donner, Pamela L.; Green, Brian E.;
ΙN
     Kempf, Dale J.; McDaniel, Keith F.; Maring, Clarence J.; Stoll, Vincent
     S.; Zhang, Rong
PΑ
     Abbott Laboratories, USA
     PCT Int. Appl., 514 pp.
     CODEN: PIXXD2
DT
     Patent
LA
     English
FAN.CNT 4
     PATENT NO.
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             LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ,
             OM, PG, PH, PL, PT, RO,
                                     RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM,
             TN, TR, TT, TZ, UA, UG}
                                     UZ, VC, VN, YU, ZA, ZM,
         RW: BW, GH, GM, KE, LS, MW;
                                     MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ,
             BY, KG, KZ, MD, RU, TJ,
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     US 2003-625121
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     WO 2003-US34707
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                                 20031031
     MARPAT 140:423711
OS
GI
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- * STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY AVAILABLE VIA OFFLINE PRINT *
- Title compds. I [wherein A = monocyclic or bicyclic ring selected from hetero/aryl, cycloalkyl, cycloalkenyl, heterocyclyl; R1 = H, (un)substituted cycloalkyl/cyclo/alkenyl, alkoxycarbonyl/alkoxy/aryl/aryls ulfonyl/arylsulfanyl/carboxy/cyano/heteroaryl/alkyl, heterocyclyl, etc.; R2, R3 = independently H, cyano, halo, (un)substituted alkenyl, alkoxycarbonyl, alkyl, heteroaryl, etc.; CR2R3C = 5- or 6-membered ring selected from Ph, pyridinyl, pyrimidinyl, pyridazinyl, thienyl, furanyl, pyrazolyl, oxazolyl, thiazolyl, imidazolyl, isoxazolyl, isothiazolyl, triazolyl, thiadiazolyl, tetrazolyl, cyclopentyl, and cyclohexyl; R4 = OH

IT

and derivs., halo, NH2 and derivs., etc.; R5 = independently CN, NO2, (un) substituted alk(en/yn)yl, hetero/aryl, arylsulfonyl, heterocyclyl etc.; n = 0-4; their pharmaceutically acceptable salts, stereoisomers, or tautomers] were prepared as hepatitis C (HCV) polymerase inhibitors for treating related infections. Thus II was prepared by alkylation of III (preparation given) with tris(methylthio)methyl Me sulfate in AcOH, cyclization with 2-amino-4[(4-methoxymethoxy)methyl]thiphene-3sulfonamide, deprotection, condensation with cyclopropanecarboxaldehyde, reduction with LiBH4. I inhibited HCV polymerase with IC50's in the range of 0.002 μM to 500 μM . I inhibited RNA replication with EC50 in the range of 0.002 μM to > 100 μM . I exhibited a cytopathic effect reduction with TC50's in the range of 6.6 μM to > 100 μM . 7481-89-2, Zalcitabine RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (combination therapy; preparation of 1,1-dioxidobenzothiadiazines as hepatitis C polymerase inhibitors and anti-infective agents) 7481-89-2 CAPLUS

RN 7481-89-2 CAPLUS CN Cytidine, 2',3'-dideoxy- (8CI, 9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

L17 ANSWER 11 OF 24 CAPLUS COPYRIGHT 2006 ACS on STN

AN 2004:332156 CAPLUS

DN 140:399402

TI Depletion of mitochondrial DNA in liver under antiretroviral therapy with didanosine, stavudine, or zalcitabine

AU Walker, Ulrich A.; Baeuerle, Jochen; Laguno, Montse; Murillas, Javier; Mauss, Stefan; Schmutz, Guenther; Setzer, Bernhard; Miquel, Rosa; Gatell, Jose M.; Mallolas, Josep

CS Department of Clinical Immunology, Medizinische Universitaetsklinik, Freiburg, Germany

SO Hepatology (Hoboken, NJ, United States) (2004), 39(2), 311-317 CODEN: HPTLD9; ISSN: 0270-9139

PB John Wiley & Sons, Inc.

DT Journal

LA English

AB The "D drug" HIV reverse-transcriptase inhibitors zalcitabine, didanosine, and stavudine are relatively strong inhibitors of polymerase-gamma compared with the "non-D drugs" zidovudine, lamivudine, and abacavir. D drugs deplete mitochondrial DNA (mtDNA) in cultured hepatocytes. This mtDNA depletion is associated with an increased in vitro production of lactate. To investigate the origin of hyperlactatemia in HIV-infected patients and the effects of antiretroviral therapy on liver mtDNA, we biopsied liver tissue from 94 individuals with chronic hepatitis C virus (HCV) infection. Eighty subjects were coinfected with HIV. Serum lactate was measured at the time of biopsy. Hepatic mtDNA and liver histol. were centrally assessed. Liver mtDNA content of HIV-infected patients receiving D drugs at the time of biopsy (n = 34) was decreased by 47% (P<.0001) compared with those without D drugs (n = 35). Aside from a possible association between HCV genotype I status and mtDNA depletion in multivariate anal., there were no other virol., immunol., histol., demog. or treatment-related variables that could explain the mtDNA depletion. Lactate was above the upper limit of normal in only three patients, all of whom were treated with D drugs. The mtDNA in each of them was lower than in any non-D drug patient and significantly (P = .017) depleted compared with D drug patients with normal lactate. In conclusion, D drug treatment is associated with decreased hepatic mtDNA in HIV-infected patients with chronic HCV infection. Moderate mtDNA depletion in liver does not necessarily lead to hyperlactatemia, but more pronounced decreases in hepatic mtDNA may be an important contributor to lactate elevation. 7481-89-2, Zalcitabine IT

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (depletion of mitochondrial DNA in liver under antiretroviral therapy

with didanosine, stavudine, or zalcitabine)
RN 7481-89-2 CAPLUS
CN Cytidine, 2',3'-dideoxy- (8CI, 9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

RE.CNT 24 THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L17 ANSWER 12 OF 24 CAPLUS COPYRIGHT 2006 ACS on STN

AN 2003:920253 CAPLUS

DN 140:350071

TI Effect of antiretroviral therapy on liver-related mortality in patients with HIV and hepatitis C virus coinfection

AU Qurishi, Nazifa; Kreuzberg, Christina; Luechters, Guido; Effenberger, Wolfgang; Kupfer, Bernd; Sauerbruch, Tilman; Rockstroh, Juergen K.; Spengler, Ulrich

CS Department of Internal Medicine, University of Bonn, Bonn, D-53105,

SO Lancet (2008), 362(9397), 1708-1713 CODEN: LANÇÃO; ISSN: 0140-6736

PB Elsevier Science Ltd.

DT Journal

LA English

Highly active antiretroviral therapy (HAART) has improved the prognosis of HIV infection. However, replication of hepatitis C virus (HCV) is not inhibited by HAART, and treatment-related hepatotoxicity is common. To clarify the effect of HAART in HIV/HCV-coinfected patients, we studied liver-related mortality and overall mortality in 285 patients who were regularly treated during the period 1990-2002 at our department. Survival was analyzed retrospectively by Kaplan-Meier and Cox's regression analyses after patients (81% hemophiliacs) had been stratified into three groups according to their antiretroviral therapy (HAART n=93, available after 1995; treatment exclusively with nucleoside analogs n=55, available after 1992; or no treatment, n=137). Liver-related mortality rates were 0.45, 0.69, and 1.70 per 100 person-years in the HAART, antiretroviral-treatment, and untreated groups. Kaplan-Meier anal. of liver-related mortality confirmed the significant survival benefit in patients with antiretroviral therapy, and regression anal. identified HAART (odds ratio 0.106 [95% CI 0.020-0.564]), antiretroviral treatment (0.283 [0.103-0.780]), CD4-pos. T-cell count (0.746 [0.641-0.868] per 0.05+109 cells/L), serum cholinesterase (0.962 [0.938-0.986] per 100 U/L), and age (1.065 [1.027-1.105] per yr) as independent predictors of liver-related survival. Severe drug-related hepatotoxicity was seen in five patients treated with nucleoside analogs alone and 13 treated with HAART. No patient died from drug-related hepatotoxicity. In addition to improved overall survival, antiretroviral therapy significantly reduced long-term liver-related mortality in our patients. This survival benefit seems to outweigh by far the associated risks of severe hepatotoxicity. 7481-89-2, Zalcitabine

RL: ADV (Adverse effect, including toxicity); PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (antiretroviral therapy effect on liver-related mortality in patients with HIV and hepatitis C virus coinfection)

RN 7481-89-2 CAPLUS

CN Cytidine, 2',3'-dideoxy- (8CI, 9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

McIntosh

IT

THERE ARE 48 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 48 ALL CITATIONS AVAILABLE IN THE RE FORMAT

L17 ANSWER 13 OF 24 CAPLUS COPYRIGHT 2006 ACS on STN

2003:772804 CAPLUS AN

DN 140:296896

- Risk of severe hepatotoxicity associated with antiretroviral therapy in the HIV-NAT Cohort, Thailand, 1996-2001
- Law, W. Phillip; Dore, Gregory J.; Duncombe, Chris J.; Mahanontharit, ΑU Apicha; Boyd, Mark A.; Ruxrungtham, Kiat; Lange, Joep M.; Phanuphak, Praphan; Cooper, David A.
- National Centre in HIV Epidemiology and Clinical Research, University of CS New South Wales, Sydney, 2010, Australia
- AIDS (London, United Kingdom) (200%), 17(15), 2191-2199 CODEN: AIDSET; ISSN: 0269-9370
- PBLippincott Williams & Wilkins
- DTJournal
- English LA
- The aim was to examine rates and predictors of severe hepatotoxicity with combination antiretroviral therapy in a developing country setting: the eight HIV-NAT randomized controlled trials in Thailand. All patients (n = 692) received at least two nucleoside reverse transcriptase inhibitors; 215 also received a non-nucleoside reverse transcriptase inhibitor (NNRTI) and 135 also received a protease inhibitor. Severe hepatotoxicity was defined as an increase in alanine aminotransferase (ALT) level to five times the upper limit of normal and an increase of at least 100 U/l from baseline. Liver function tests were available at baseline and weeks 4, 8, 12, 24, 36 and 48. Hepatitis B virus (HBV) and hepatitis C virus (HCV) testing was performed on stored serum. Mean age was 32.3 yr; 52% were male, 11% had Centers for Disease Control and Prevention category C HIV disease at baseline, and 22% had received prior antiretroviral therapy. Prevalence of HBV, HCV and HBV/HCV coinfection was 8.7%, 7.2%, and 0.4%, resp. Incidence of severe hepatotoxicity was 6.1/100 person-years [95% confidence interval (CI), 4.3-8.3/100]. In multivariate anal., predictors of severe hepatotoxicity were HBV or HCV coinfection, and NNRTI-containing therapy. Incidence of severe hepatotoxicity was particularly high among patients receiving nevirapine (18.5/100 person-years; 95% CI, 11.6-27.8) and nevirapine/efavirenz (44.4/100 person-years; 95% CI, 12.1-113.7). Incidence and risk factors for severe hepatotoxicity appear similar among these Thai patients to those in other racial groups. Development of standardized antiretroviral therapy regimens for developing country settings should consider potential toxicity and capabilities for monitoring of toxicity.
- 7481-89-2, Zalcitabine

RL: ADV (Adverse effect, including toxicity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(risk of severe hepatotoxicity associated with antiretroviral therapy in HIV-infected patients)

7481-89-2 CAPLUS RN

Cytidine, 2',3'-dideoxy- (8CI, 9CI) (CA INDEX NAME) CN

Absolute stereochemistry. Rotation (+).

RE.CNT 23 THERE ARE 23 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L17 ANSWER 14 OF 24 CAPLUS COPYRIGHT 2006 ACS on STN
- 2003:347498 CAPLUS
- 139:47738 DN
- Performance characteristics of the TRUGENE HIV-1 genotyping kit and the OpenGene DNA sequencing system
- Kuritzkes, Daniel R.; Grant, Robert M.; Feorino, Paul; Griswold, Marshal; AU Hoover, Marie; Young, Russell; Day, Stephen; Lloyd, Robert M., Jr.; Reid, Caroline; Morgan, Gillian F.; Winslow, Dean L.
- CS Division of Infectious Diseases, University of Colorado Health Sciences

Center, Denver, CO, USA Journal of Clinical Microbiology (2003), 41(4), 1594-1599 CODEN: JCMIDW; ISSN: 0095-1137

PBAmerican Society for Microbiology

 \mathtt{DT} Journal English LA

The TRUGENE HIV-1 Genotyping Kit and OpenGene DNA Sequencing System are designed to sequence the protease (PR) - and reverse transcriptase (RT)-coding regions of human immunodeficiency virus type 1 (HIV-1) pol. Studies were undertaken to determine the accuracy of this assay system in detecting resistance-associated mutations and to determine the effects of RNA extraction methods, anticoagulants, specimen handling, and potentially interfering substances. Samples were plasma obtained from HIV-infected subjects or seroneg. plasma to which viruses derived from wild-type and mutant infectious mol. clones (IMC) of HIV-1 were added. Extraction methods tested included standard and UltraSensitive AMPLICOR HIV-1 MONITOR, QIAGEN viral RNA extraction mini kit, and QIAGEN Ultra HIV extraction kit, and NASBA manual HIV-1 quant. NucliSens. Sequence data from test sites were compared to a "gold standard" reference sequence to determine the percent agreement. Comparisons between test and reference sequences at the nucleotide level showed 97.5 to 100% agreement. Similar results were obtained regardless of extraction method, regardless of use of EDTA or acid citrate dextrose as anticoagulant, and despite the presence of triglycerides, bilirubin, Hb, antiretroviral drugs, HIV-2, hepatitis C virus (HCV), HBV, cytomegalovirus, human T-cell leukemia virus type 1 (HTLV-1), or HTLV-2. Samples with HIV-1 RNA titers of ≥1,000 copies/mL gave consistent results. The TRUGENE HIV-1 Genotyping Kit and OpenGene DNA Sequencing System consistently generate highly accurate sequence data when tested with IMC-derived HIV and patient samples.

7481-89-2, Zalcitabine

RL: BSU (Biological study, unclassified); BIOL (Biological study) (potentially interfering substances have no impact on performance characteristics of TRUGENE HIV-1 genotyping kit and OpenGene DNA sequencing system)

7481-89-2 CAPLUS RN

Cytidine, 2',3'-dideoxy- (8CI, 9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

RE.CNT 17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L17 ANSWER 15 OF 24 CAPLUS COPYRIGHT 2006 ACS on STN

2003:26945 CAPLUS AN

139:381 DN

Ribonucleoside analogue that blocks replication of bovine viral diarrhea and hepatitis C viruses in culture

Stuyver, Lieven J.; Whitaker, Tony; McBrayer, Tamara R.; Hernandez-Santiago, Brenda I.; Lostia, Stefania; Tharnish, Phillip M.; Ramesh, Mangala; Chu, Chung K.; Jordan, Robert; Shi, Junxing; Rachakonda, Suguna; Watanabe, Kyoichi A.; Otto, Michael J.; Schinazi, Raymond F.

CS

Pharmasset Inc., Tucker, GA, 30084, USA Antimicrobial Agents and Chemotherapy (2003), 47(1), 244-254 CODEN: AMACCQ; ISSN: 0066-4804

PBAmerican Society for Microbiology

 \mathtt{DT} Journal

English LA

AB A base-modified nucleoside analog, β -D-N4-hydroxycytidine (NHC), was found to have antipestivirus and antihepacivirus activities. This compound inhibited the production of cytopathic bovine viral diarrhea virus (BVDV) RNA in a dose-dependent manner with a 90% effective concentration (EC90) of 5.4 μ M, an observation that was confirmed by virus yield assays (EC90 = 2 μ M). When tested for hepatitis C virus (HCV) replicon RNA reduction in Huh7 cells, NHC had an EC90 of 5 μ M on day 4. The HCV RNA reduction was incubation time and nucleoside concentration dependent. The in vitro antiviral effect of NHC was additive with recombinant alpha

 \mathbf{IT}

interferon-2a and could be prevented by the addition of exogenous cytidine and uridine but not of other natural ribo- or 2'-deoxynucleosides. When HCV RNA replicon cells were cultured in the presence of increasing concns. of NHC (up to 40 μM) for up to 45 cell passages, no resistant replicon was selected. Similarly, resistant BVDV could not be selected after 20 passages. NHC was phosphorylated to the triphosphate form in Huh7 cells, but in cell-free HCV NS5B assays, synthetic NHC-triphosphate (NHC-TP) did not inhibit the polymerization reaction. Instead, NHC-TP appeared to serve as a weak alternative substrate for the viral polymerase, thereby changing the mobility of the product in polyacrylamide electrophoresis gels. We speculate that incorporated nucleoside analogs with the capacity of changing the thermodn. of regulatory secondary structures (with or without introducing mutations) may represent an important class of new antiviral agents for the treatment of RNA virus infections, especially HCV. 3258-02-4, N4-Hydroxycytidine RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(N4-hydroxycytidine blocks replication of bovine viral diarrhea and hepatitis C viruses in culture)

RN 3258-02-4 CAPLUS
CN Uridine, 4-oxime (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RE.CNT 47 THERE ARE 47 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

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L17 ANSWER 16 OF 24 CAPLUS COPYRIGHT 2006 ACS on STN
     2002:927626 CAPLUS
ΑN
     138:20431
DN
     Use of mitochondrial DNA-specific quantitative real-time PCR for diagnosis
     and monitoring drug toxicity in humans suffering with various disorders
     such as viral infections, neurological disorders, cancer, arthritis, male
     sterility or organ failure
IN
     Cote, Helene; Montaner, Julio; O'Shaughnessy, Michael V.
     The University of British Columbia, Can.
PA
SO
     PCT Int. Appl., 37 pp.
     CODEN: PIXXD2
DT
     Patent
     English
LA
FAN.CNT 1
     PATENT NO.
                                             APPLICATION NO.
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                                 2002 1 205
     WO 2002097124
                          A1
                                             WO 2002-CA796
                                                                     20020529
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         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
             LS, LT, LU, LV, MA, MD;
                                     MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH,
             PL, PT, RO, RU, SD, SE,
                                     SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ,
             UA, UG, US, UZ, VN, YU,
                                     ZA, ZM, ZW
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH,
             CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR,
             BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
     CA 2416332
                          A1
                                 20021205
                                             CA 2002-2416332
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     US 2003099933
                          A1
                                 20030529
                                             US 2002-158543
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     EP 1395681
                          A1
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                                             EP 2002-729732
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20060726

20041021

20060815

20010529

20020529

IE, SI, LT, LV, FI, RO, MK, CY, AL, TR

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,

JP 2003-500289

AT 2002-729732

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McIntosh

EP 1395681

AT 334232

PRAI US 2001-293523P

JP 2004532043

WO 2002-CA796

The invention discloses the use of quant. real-time polymerase chain reaction (PCR) to monitor drug toxicity, which involves measuring the relative amount of mitochondrial DNA in peripheral blood cells obtained from individuals suffering with various disorders. The invention relates that the quant. real-time PCR involves co-amplification of a mitochondrial sequence and a reference sequence, such as a genomic sequence. The invention also discloses that said disorders include HIV infection, cancer, hepatitis A, hepatitis B, hepatitis C, arthritis, Alzheimer's disease, Parkinson's disease, or Huntington's disease. The invention also relates that said drugs used to treat patients include nucleoside or nucleotide analogs, and/or reverse transcriptase inhibitors. The invention further discloses that the said method can be used to diagnose conditions such as male infertility and organ failure. The method was illustrated by detecting the amount of mitochondrial gene CCOI and the nuclear gene ASPOLy in HIV infected individuals undergoing antiviral therapy. 7481-89-2, Hivid IT

RL: ADV (Adverse effect, including toxicity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(mitochondrial DNA-specific quant. real-time PCR for monitoring drug toxicity in individuals suffering for various disorders such as viral infections, neurol. disorders, cancer, and arthritis)

RN 7481-89-2 CAPLUS

CN Cytidine, 2',3'-dideoxy- (8CI, 9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

RE.CNT 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

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L17 ANSWER 17 OF 24 CAPLUS COPYRIGHT 2006 ACS on STN
AN
     2002:869219 CAPLUS
    137:363028
DN
     Drug screening assays and kits for discovery of anti-microbial and
TI
     chemotherapeutics agents
    McCarthy, Lawrence; Kong, Lilly; Shao, Tang; Su, Xin
     Focus Technologies, Inc., USA
     PCT Int. Appl., 94 pp.
     CODEN: PIXXD2
DT
     Patent
LΑ
     English
FAN.CNT 1
     PATENT NO.
                         KIND
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WO 2002090993
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                                             WO 2001-US44783
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     WO 2002090993
                          Α3
                                20040415
         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
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             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
             LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH,
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             KG, KZ, MD, RU, TJ, TM, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB,
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     EP 1435000
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                                             EP 2001-273944
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
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PRAI US 2000-253150P
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    US 2001-304533P
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    US 2001-297686P
                          P
                                20010712
    US 2001-996187
                          A2
                                20011127
    WO 2001-US44783
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                                20011127
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AB Methods and compns. for detecting the phenotype of a bioactive mol. assays. More specifically, are provided methods and compns. are provided for determining the suitability of one ore more candidate compds. prior to or during the course of chemotherapy or anti-infective therapy, for their capacity to inhibit the bioactive mols. of micro-organisms, cancers and as an assay for expression in transgene therapy. Also provided are phenotypic assays for drug discovery. Claimed sequences were not present at the time of publication.

IT 7481-89-2, Zalcitabine

RL: BSU (Biological study, unclassified); BIOL (Biological study) (drug screening assays for discovery of anti-microbial and chemotherapeutics agents)

RN 7481-89-2 CAPLUS

CN Cytidine, 2',3'-dideoxy- (8CI, 9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (*).

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L17 ANSWER 18 OF 24 CAPLUS COPYRIGHT 2006 ACS on STN
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AN 2002:314958 CAPLUS

DN 136:340939

TI Preparation of modified nucleosides for treatment of viral infections and abnormal cellular proliferation

IN Stuyver, Lieven; Watanabe, Kyoichi A.

PA Pharmasset Limited, USA

SO PCT Int. Appl., 230 pp.

CODEN: PIXXD2

DT Patent

LA English

Parent

FAN.	CNT 2																	
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			CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,
			HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	LK,	LR,	LS,
			LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NO,	NZ,	PL,	PT,	RO,
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$$R^{1}$$
 R^{1}
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 R^{3}
 R^{2}
 R^{2}
 R^{2}

Modified nucleosides, e.g. I, wherein D is hydrogen, alkyl, acyl, monophosphate, diphosphate, triphosphate, monophosphate ester, diphosphate ester, triphosphate ester, phospholipid or amino acid; X is H, halogen, NH2, substituted amine, oxime, OH, alkoxy, SH, thioalkyl; Y is O, S, Se; R and R1 are independently H, alkyl, alkenyl, alkynyl, aryl, alkylaryl, halogen, NH2, substituted amine, oxime, hydrazine, OH, alkoxy, SH, thioalkyl, NO2, NO, CH2OH, CH2OH, ester, CONH2, amide, CN; R2 and R3 are independently H, halogen, OH, SH, OMe, SMe, NH2, NHMe, CH:CH2, CN, CH2NH2, CH2OH, CO2H; were prepared for treating a Flaviviridae (including BVDV and HCV), Orthomyxoviridae (including Influenza A and B) or Paramyxoviridae (including RSV) infection, or conditions related to abnormal cellular proliferation, in a host, including animals, and especially humans. This invention also provides an effective process to quantify the viral load, and in particular BVDV, HCV or West Nile Virus load, in a host, using real-time polymerase chain reaction ("TR-PCR"). Addnl., the invention discloses probe mols. that can fluoresce proportionally to the amount of virus present in a sample. Thus, (1'R, 2'S, 3'R, 4'R)-1-[2, 3-1]dihydroxy-4-(hydroxymethyl)cyclopentan-1-yl}-5-fluorocytosine was prepared and tested in vitro as antiviral and antitumor agent. 3258-02-4P ΙT

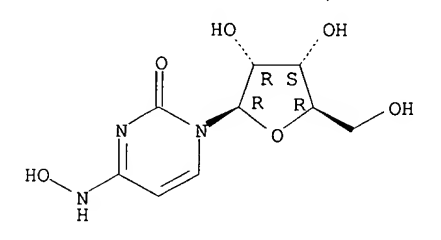
RL: IMF (Industrial manufacture); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of modified nucleosides for treatment of viral infections and

abnormal cellular proliferation)

RN 3258-02-4 CAPLUS
CN Uridine, 4-oxime (9CI) (CA INDEX NAME)

Absolute stereochemistry.



L17 ANSWER 19 OF 24 CAPLUS COPYRIGHT 2006 ACS on STN

AN 2002:171918 CAPLUS

DN 136:217007

TI Preparation of antiviral nucleoside derivatives as inhibitors of subgenomic hepatitis C virus RNA replication

IN Devos, Rene; Dymock, Brian William; Hobbs, Christopher John; Jiang,
Wen-rong; Martin, Joseph Armstrong; Merrett, John Herbert; Najera, Isabel;
Shimma, Nobuo; Tsukuda, Takuo

PA F. Hoffmann-La Roche Ag, Switz.

SO PCT Int. Appl., 225 pp.

CODEN: PIXXD2

DT Patent

LA English FAN CNT 1

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	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE		
ΡI	WO 2002018404	A2	20020307	WO 2001-EP9633	20010821		
	WO 2002018404	A9	20031002				

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    MARPAT 136:217007
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HO
$$X \rightarrow B$$
 $A \rightarrow B$ A

- Nucleosides I , wherein Rl is hydrogen, hydroxy, alkyl, hydroxyalkyl, alkoxy, halogen, cyano, isocyano or azido; R2 is hydrogen, hydroxy, alkoxy, chlorine, bromine or iodine; R3 is hydrogen; or R2 and R3 together represent =CH2; or R2 and R3 represent fluorine; X is O, S or CH2; B is a substituted purine base, were prepared as inhibitors of subgenomic hepatitis C virus (HCV) RNA replication. Thus, nucleoside II was prepared and tested for the inhibition of HCV RNA replication (EC50 = 0.6 μM).
- 3258-02-4P 7481-89-2P ITRL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

(preparation of antiviral nucleoside derivs. as inhibitors of subgenomic Crds 239-241 hepatitis C virus RNA replication)

RN 3258-02-4 CAPLUS

Uridine, 4-oxime (9CI) (CA INDEX NAME) CN

Absolute stereochemistry.

7481-89-2 CAPLUS RN

CN Cytidine, 2',3'-dideoxy- (8CI, 9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

L17 ANSWER 20 OF 24 CAPLUS COPYRIGHT 2006 ACS on STN

AN 2002:109650 CAPLUS

DN 136:288583

TI Effects of HAART on hepatitis C, hepatitis G, and TT virus in multiply coinfected HIV-positive patients with haemophilia

AU Takamatsu, J.; Toyoda, H.; Fukuda, Y.; Nakano, I.; Yokozaki, S.; Hayashi, K.; Saito, H.

CS Department of Transfusion Medicine, Nagoya University School of Medicine, Nagoya, 466-8550, Japan

SO Haemophilia (2001), 7(6), 575-581 CODEN: HAEMF4; ISSN: 1351-8216

PB Blackwell Science Ltd.

DT Journal

LA English

In multiply coinfected human immunodeficiency virus (HIV)-pos. patients, we investigated the effects of high-activity antiretroviral therapy (HAART) using HIV protease inhibitors on three other viruses: hepatitis C virus (HCV), hepatitis G virus (HGV), and TT virus (TTV). Viral concns. were measured serially by polymerase chain reaction methods in five patients with quadruple infection (HIV, HCV, HGV, and TTV) and in two patients with triple infection (HIV, HCV, and HGV) before and during HAART. In addition, CD4+ cell counts and serum alanine aminotransferase (ALT) levels were measured serially. Generally we observed no difference in serum HCV RNA, HGV RNA, or TTV DNA concns. between samples obtained before and after initiation of HAART, whereas HIV RNA concentration decreased and CD4 counts increased in most patients. However, two patients had markedly decreased concns. of HCV RNA and HGV RNA, resp., more than 12 mo after beginning HAART. Normalization of serum ALT levels was observed in a patient with decline of HCV RNA concns. No interactions were observed among these four viruses. HAART had no apparent direct effects on HCV, HGV, or TTV. Further studies will be required to elucidate whether the restoration of immune status through suppression of HIV replication by HAART may affect HCV or HGV RNA concns.

IT 7481-89-2, Zalcitabine

RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(HAART effect on hepatitis C, hepatitis G, and TT virus in HIV-pos. patients with multiple coinfections and haemophilia)

RN 7481-89-2 CAPLUS

CN Cytidine, 2',3'-dideoxy- (8CI, 9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

RE.CNT 49 THERE ARE 49 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L17 ANSWER 21 OF 24 CAPLUS COPYRIGHT 2006 ACS on STN

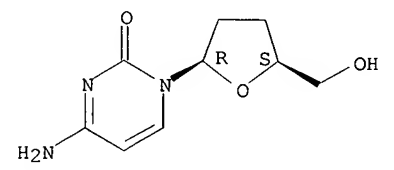
AN 2002:107667 CAPLUS

DN 136:145568

TI Improved tolerance to anti-viral and anti-tumor chemotherapy by administration of erythropoietin

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IN
    Itri, Loretta; Bowers, Peter
    Ortho-McNeil Pharmaceutical, Inc., USA
PA
    PCT Int. Appl., 56 pp.
     CODEN: PIXXD2
DT
     Patent
     English
LA
FAN.CNT 1
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PRAI US 2000-222538P
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    WO 2001-US24426
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    The present invention provides methods using erythropoietin to improve the
     tolerance of anti-viral and anti-tumor chemotherapeutic regimens containing
                 The invention also described improved methods to treat
    chronic HCV by adjusting the dose of ribavirin to tailor the
     active dose of the drug while supporting the Hb levels in the patient with
     EPO. The present invention also provides anti-viral dosing regimens,
     particularly for chronic HCV comprising administration of an
     interferon containing anti-viral medicament, EPO, and a compound that reduces
     the amount of active tumor necrosis factor in the subject.
    7481-89-2, Zalcitabine
IT
    RL: ADV (Adverse effect, including toxicity); PAC (Pharmacological
     activity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
        (improved tolerance to anti-viral and anti-tumor chemotherapy by
        administration of erythropoietin)
   7481-89-2 CAPLUS
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Absolute stereochemistry. Rotation (+).



RE.CNT 14 THERE ARE 14 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

Cytidine, 2',3'-dideoxy- (8CI, 9CI) (CA INDEX NAME)

L17 ANSWER 22 OF 24 CAPLUS COPYRIGHT 2006 ACS on STN

AN 2001:784185 CAPLUS

DN 136:95621

TI Low frequency of severe hepatotoxicity and association with HCV coinfection in HIV-positive patients treated with HAART

AU Monforte, Antonell d'Arminio; Bugarini, Roberto; Pezzotti, Patrizio; De Luca, Andrea; Antinori, Andrea; Mussini, Cristina; Vigevani, Gian Marco; Tirelli, Umberto; Bruno, Raffaele; Gritti, Francesco; Piazza, Marcello; Chigiotti, Silvia; Chirianni, Antonio; De Stefano, Carlo; Pizzigallo, Eligio; Perrella, Oreste; Moroni, Mauro

CS ICONA Study Group, Institute of Infectious and Tropical Diseases, L Sacco H, University of Milan, Milan, 20157, Italy

JAIDS, Journal of Acquired Immune Deficiency Syndromes (2001), 28(2),

McIntosh

114-123

CN

CODEN: JJASFJ

PB Lippincott Williams & Wilkins

DT Journal

LA English

Highly active antiretroviral therapy (HAART) is strongly effective in reducing morbidity and mortality in HIV-1-pos. individuals. Its main drawback is the potential toxicity. Data on the frequency and determinants of severe hepatotoxicity in a clin. setting are still sparse. This is a prospective study of HIV-1-pos. individuals with known HBsAg and HCV-Ab serol. The study end point was progression to alanine aminotransferase (ALT) levels ≥200 IU/L after HAART initiation. Cumulative probability of progression to this end point was estimated by the Kaplan-Meier method. Crude and adjusted hazard ratios (HR) were estimated by proportional hazards regression model. One thousand two hundred fifty-five patients were included. HBsAg was found in 91 (7.2%), HCV-Ab in 578 (46.5%) patients; almost all injection drug users (451 of 482; 93.6%) were HCV-Ab pos. Sixty-one individuals progressed to the end point with a probability of 7.9% (95% confidence interval [CI], 5.6-10.0) of progression at 24 mo from starting. Independent factors predicting progression to the end point were baseline ALT levels (HR, 5.29; 95% CI, 3.24-8.65; every 10 IU/L higher), HCV-Ab positivity (HR, 4.01; 95% CI, 1.48-10.85) or both HBsAg and HCV-Ab positivity (HR, 3.85, 95% CI, 1.01-14.61), and previous non-HAART therapy (HR, 1.84, 95% CI, 1.04-3.42). Patients receiving stavudine-containing regimens had a lower risk than those receiving zidovudine-containing regimens (HR, 0.30, 95% CI, 0.12-0.71). There was a low risk of ALT ≥200 IU/L in the authors' cohort. Hepatitis C coinfection and elevated ALT levels at HAART initiation are important predictors of progression to ALT ≥200 IU/L; stavudine-containing regimens were associated with a lower risk compared with zidovudine-containing regimens.

IT 7481-89-2, Zalcitabine

RL: ADV (Adverse effect, including toxicity); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(low frequency of severe hepatotoxicity and association with HCV coinfection in HIV-pos. humans treated with HAART)

RN 7481-89-2 CAPLUS

CN Cytidine, 2',3'-dideoxy- (8CI, 9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

RE.CNT 24 THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L17 ANSWER 23 OF 24 CAPLUS COPYRIGHT 2006 ACS on STN

AN 2000:840382 CAPLUS

DN 135:40464

TI Safety and efficacy of interferon-ribavirin combination therapy in HCV-HIV coinfected subjects: An early report

AU Zylberberg, H.; Benhamou, Y.; Lagneaux, J. L.; Landau, A.; Chaix, M. -L.; Fontaine, H.; Bochet, M.; Poynard, T.; Katlama, C.; Pialoux, G.; Brechot, C.; Pol, S.

CS Unite d'Hepatologie, INSERM U370, Unite d'Hepatologie, INSERM U370, CHU Necker, Paris, Fr.

SO Gut (2000), 47(5), 694-697

CODEN: GUTTAK; ISSN: 0017-5749
PB BMJ Publishing Group

DT Journal

LA English

More severe liver disease together with a poor response rate to α interferon argue for the use of more potent anti-hepatitis C virus (HCV) therapies in human immunodeficiency virus (HIV)-HCV coinfected patients, but the efficacy and safety of interferon-ribavirin combination therapy in HIV infected subjects are unknown. Aim of this study was to retrospectively evaluate the efficacy and safety of anti-HCV combination therapy in 21 HCV-HIV coinfected

patients receiving antiretroviral therapy, and to access the clin. relevance of in vitro inhibition of phosphorylation by ribavirin of potent inhibitors of HIV-i.e., zidovudine, stavudine, and zalcitabine. Twenty one patients were treated with combined antiretroviral therapy including zidovudine (n=8) or stavudine (n=13) (in association with protease inhibitors in 12). All received ribavirin (1000 or 1200 mg/day) and α interferon (3 MU three times/wk) for chronic hepatitis C infection. All patients had not responded (n=20) or relapsed (n=1) after a previous six month course of α interferon therapy. HIV viral load (Monitor test) and CD4 cells count were measured at the beginning and every three months during and after ribavirin plus α interferon therapy over a mean period of 11 (1) months. Clin. and biol. adverse effects were recorded. There was no significant variation in HIV viral load or CD4 cell counts after three or six months of ribavirin therapy compared with baseline values. Of the 21 subjects, three (14%) had an increase in HIV viral load of more than 0.5 log leading to discontinuation of ribavirin in one. Eleven of 21 (52.4%) and initial neg. HCV viremia at three (n=10) or six (n=1) months but only six were polymerase chain reaction neg. at the end of therapy, leading to rates for primary response and breakthrough of 23.8% and 28.5%, resp. Six months after completion of therapy, three patients relapsed (14.3%) and three (14.3%) had sustained virol. response. Median Hb concentration decreased significantly after three and six months of ribavirin therapy (p=0.0002 and p=0.0003, resp.) leading to withdrawal of therapy in one patient. These preliminary results show that: (1) despite in vitro interactions between ribavirin, zidovudine, and stavudine, significant variation in HIV replication does not usually occur in HCV-HIV coinfected patients receiving ribavirin and different antiretroviral regimens, including zidovudine and stavudine; (2) α interferon and ribavirin combination therapy induced primary and sustained virol. responses in 28.5% and 14.3% of treated subjects (who were previous non-responders to interferon therapy), resp.; (3) anemia is a frequent adverse event. Such results should be confirmed in larger prospective trials.

IT 7481-89-2, Zalcitabine

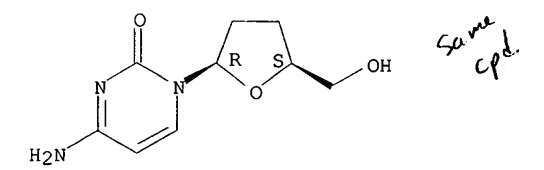
RL: ADV (Adverse effect, including toxicity); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(interferon- α and ribavirin combination therapy in humans coinfected with hepatitis C virus and HIV)

RN 7481-89-2 CAPLUS

CN Cytidine, 2',3'-dideoxy- (8CI, 9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).



RE.CNT 13 THERE ARE 13 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L17 ANSWER 24 OF 24 CAPLUS COPYRIGHT 2006 ACS on STN

AN 2000:443717 CAPLUS

DN 133:37763

TI Can HCV affect the efficacy of anti-HIV treatment?

AU Filippini, P.; Coppola, N.; Scolastico, C.; Liorre, G.; Nocera, R.; Sagnelli, E.; Piccinino, F.

CS Institute of Infectious Diseases, School of Medicine, Second University of Naples, Naples, Italy

SO Archives of Virology (2000), 145(5), 937-944 CODEN: ARVIDF; ISSN: 0304-8608

PB Springer-Verlag Wien

DT Journal

LA English

AB To evaluate the impact of new antiretroviral combinations (HAART: Highly Active Anti Retroviral Therapy) on HCV replication and liver enzyme levels, we analyzed the changes in HCV viremia and aminotransferase levels in HIV and HCV co-infected patients.

Moreover, to evaluate the influence of HCV infection on the efficacy of HAART, we compared the virol., immunol. and biochem. response

IT

to antiretroviral combinations in anti-HIV pos. subjects with or without HCV infection. We enrolled eight consecutive outpatients with HIV-HCV coinfection and with indications for HAART (Group A). For each patient in group A, we selected an anti-HIV neg. patient with indications for HAART, pair-matched for age, sex, risk factor for HIV infection, presumed duration of infection, number of CD4 cells, HIV viremia and treatment schedule (Group B). A statistically significant increase in CD4 in both groups was found at 1st, 3rd and 6th month of antiretroviral therapy. A decrease in HIV-RNA in both groups was observed at 1st and 6th month of treatment. The percentage of patients with undetectable HIV-RNA at the 1st month was higher in Group B than in Group A (8/8 vs. 3/8, p = 0.025). Basal HCV-RNA viremia was very high in each case and no variations during treatment were observed During therapy the aminotransferase levels slightly decreased in Group A and consistently increased in Group B. In Group A the differences were not significant to the statistical anal.; in Group B the aminotransferase levels at 3rd and 6th month were significantly higher than those observed at the baseline. 7481-89-2, Zalcitabine RL: ADV (Adverse effect, including toxicity); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (can HCV affect efficacy of anti-HIV treatment)

RN 7481-89-2 CAPLUS

CN Cytidine, 2',3'-dideoxy- (8CI, 9CI) (CA INDEX NAME)

Absolute stereochemistry. Rotation (+).

RE.CNT 25 THERE ARE 25 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

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